# **WEST Search History**



DATE: Wednesday, March 02, 2005

Hide?	Set Name	Query	<u>Hit</u> Count			
	DB=PGPB; PLUR=YES; OP=ADJ					
Ü	L37	L32 and (stability)	1			
	L36	L32 and (stable)	0.			
	L35	L32 and (stable adj2 air)	0			
	L34	L32 and (absence adj2 solvent)	0			
	L33	L32 and quantum yield	1			
	L32	20030178607	1			
	DB=USPT; PLUR=YES; OP=ADJ					
	L31	6783814.pn.	1			
	DB = PGPB, USPT, USOC, EPAB, JPAB, DWPI, TDBD; PLUR = YES; OP = ADJ					
	L30	(persistence adj2 nm)	1			
	L29	polymer with (persistence adj2 nm)	0			
	L28	L26 and iptycene	6			
	L27	L26 and fluorescen\$3	10			
	L26	L25 or 124 or 123	132			
	L25	zhu-zhengguo\$.in.	4			
	L24	long-timothy\$.in.	109			
	L23	swager-timothy\$.in.	. 25			
	DB=	DB=USPT,EPAB,JPAB,DWPI; PLUR=YES; OP=ADJ				
		121 and fluorescence with yield	4			
	DB=	PGPB, USPT; PLUR=YES; OP=ADJ				
	L21  DB=	US-4356429-\$.DID. OR US-4687732-\$.DID. OR US-4927768-\$.DID. OR US-4946890-\$.DID. OR US-4992302-\$.DID. OR US-5155149-\$.DID. OR US-5194393-\$.DID. OR US-5236808-\$.DID. OR US-5244813-\$.DID. OR US-5254633-\$.DID. OR US-5364797-\$.DID. OR US-5414069-\$.DID. OR US-5451683-\$.DID. OR US-0551547-\$.DID. OR US-5512490-\$.DID. OR US-5532129-\$.DID. OR US-5540999-\$.DID. OR US-5546889-\$.DID. OR US-554747-\$.DID. OR US-5556524-\$.DID. OR US-5563056-\$.DID. OR US-556322-\$.DID. OR US-5580527-\$.DID. OR US-5585646-\$.DID. OR US-5591787-\$.DID. OR US-5597890-\$.DID. OR US-5607864-\$.DID. OR US-5679773-\$.DID. OR US-5700696-\$.DID. OR US-5705348-\$.DID. OR US-5709994-\$.DID. OR US-5710197-\$.DID. OR US-5723218-\$.DID. OR US-5869562-\$.DID. OR US-6020426-\$.DID. OR US-6259277-\$.DID. OR US-6920426-\$.DID. OR US-6259277-\$.DID.	36			

L20	L19 and alkoxyamine	5
L19	L18 orl17 or l16	665
L18	ujikawa-norihisa\$.in.	5
L17	nakamura-tomoyuki\$.in.	192
L16	hayashi-masaki\$.in.	665
DB =	USPT,USOC; PLUR=YES; OP=ADJ	
L15	3969071.pn.	1
DB =	PGPB,USPT,USOC,EPAB,JPAB,DWPI,TDBD; PLUR=YES; OP=ADJ	
L14	(alkoxybenzene or methoxybenzene)(sulfonic) with (polymer\$7 or poly or polycondensation or condensation or \$3ion exchange)	5
L13	alkoxybenzene sulfonic with (polymer\$7 or poly or polycondensation or condensation)	0
L12	L10 not 19	1
L11	L'10 not 18	0
L10	L8 and (poly or polymer\$7 or polycondensation)	12
L9	L8 and (polymer\$7 or polycondensation)	11
L8	(sulfonated or sulfated or sulfonic) adj3 (\$2methoxy benzene or \$2methoxybenzene)	19
L7	(sulfonated or sulfated or sulfonic) adj (\$2methoxy benzene or \$2methoxybenzene)	1
L6	(sulfonated or sulfated or sulfonic) with methoxy benzene	181
L5	12 same (sulfonated or sulfated or sulfonic)	6
L4	12 same (sulfonated or sulfated or sulfo)	0
L3	(sulfonated or sulfated or sulfo) with L2	0
L2	(polymer&7 or poly\$5)with (DMB or dimethoxybenzene or dialkoxybenzene or dibutoxybenzene)	181
L1	(polymer&7 or poly\$5)adj2 (DMB or dimethoxybenzene or dialkoxybenzene or dibbutoxybenzene)	9

# **END OF SEARCH HISTORY**

=> file reg FILE 'REGISTRY' USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 American Chemical Society (ACS)

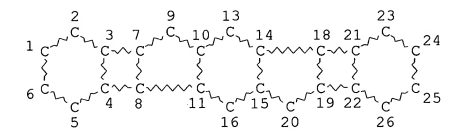
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FILE 'HCAPLUS'
L1
          368 S SWAGER ?/AU
          2660 S ZAHN ?/AU
L2
             2 S L1 AND L2
L3
               SEL L3 1-2 RN
    FILE 'REGISTRY'
          2 S E1-E2
L4
    FILE 'LREGISTRY'
L5
               STR
L6
               STR
    FILE 'REGISTRY'
L7
            0 S L5 AND L6
L8
             0 S L5
L9
             STR
L10
             0 S L9
             2 S L9 FUL
L11
               SAV L11 ZEM041/A
    FILE 'CAOLD'
          0 S L11
L12
    FILE 'ZCAPLUS'
L13
            1 S L11
   FILE 'REGISTRY'
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STR

=> d l11 que stat

L9



NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 24

STEREO ATTRIBUTES: NONE

L11 2 SEA FILE=REGISTRY SSS FUL L9

100.0% PROCESSED 382715 ITERATIONS

SEARCH TIME: 00.00.07

2 ANSWERS

=> file zcaplus FILE 'ZCAPLUS' USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2005 AMERICAN CHEMICAL SOCIETY (ACS)

### => d 113 1 all hitstr

L13 ANSWER 1 OF 1 ZCAPLUS COPYRIGHT 2005 ACS on STN

AN 2000:619390 ZCAPLUS

DN 134:21651

ED Entered STN: 06 Sep 2000

TI Special geminals and Schlegel diagrams of molecular structures of fullerenes and metallofullerenes

AU Chiu, Y.-N.; Xiao, J.; Merritt, C. D.; Liu, K.; Huang, W.-X.; Ganelin, P. V.; Li, N. N.

CS Department of Chemistry, Center for Molecular Dynamics and Energy Transfer, The Catholic University of America, Washington, DC, 20064, USA

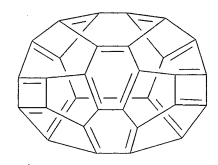
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THEOCHEM (2000), 530(1,2), 67-83
SO
     CODEN: THEODJ; ISSN: 0166-1280
PB
     Elsevier Science B.V.
DT
     Journal
     English
LA
     65-3 (General Physical Chemistry)
CC
     Schlegel diagrams were used to demonstrate the location of geminals
AB
     for the following mol. systems: C42H28 .fwdarw. C42 (C2v, D2h),
     C30H18 .fwdarw. C30 (C2v, D5h), C4H4 (D4h), C4H6, C28 (Td), Ti@C28
     (Td), Sc3@C82 (C3v), and Sc@C20 (D5).
     geminal Schlegel diagram fullerene metallofullerene
ST
ΙT
     Wave function
        (geminal; geminals and Schlegel diagrams of fullerenes and
        metallofullerenes)
    Molecular orbital
IT
    Molecular structure-property relationship
    Molecular topology
        (geminals and Schlegel diagrams of fullerenes and
        metallofullerenes)
ΙT
     Fullerenes
     Fullerides
        (geminals and Schlegel diagrams of fullerenes and
        metallofullerenes)
IT
     106-99-0, 1,3-Butadiene, properties
                                           517-51-1
                                                      1120-53-2,
     1,3-Cyclobutadiene 10075-85-1 115383-19-2, [5,6]Fullerene-C28-Td
     135026-72-1, [5,6]Fullerene-C30-D5h 145077-51-6
                                                         146750-44-9
     309242-94-2, [5,6,7] Fullerene-C42-C2v 309242-95-3,
     [4,5,6,7]Fullerene-C42-D2h 309242-96-4, [4,5,6]Fullerene-C30-C2v
     309242-97-5, [4,5,6,7] Fullerene-C42-C2v 309242-98-6,
     [4,5,6]Fullerene-C30-C2v 309244-06-2
        (geminals and Schlegel diagrams of fullerenes and
        metallofullerenes)
RE.CNT
              THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS RECORD
        21
RE
(1) Baum, R; Chem Engng News 1988, V29, P33
(2) Chiu, Y; Acta Phys Hungarica 1994, V74(4), P427 ZCAPLUS
(3) Chiu, Y; Chem J Chin Univ 1997, V18, P1147 ZCAPLUS
(4) Chiu, Y; Eur J Solid State Inorg Chem 1993, Vt30, P1119
(5) Chiu, Y; J Chin Chem Soc 1992, V37, P361
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(7) Chiu, Y; J Mol Struct (Theochem) 1994, V312, P215
(8) Chiu, Y; J Mol Struct (Theochem) 1995, V332, P47 ZCAPLUS
(9) Chiu, Y; J Mol Struct (Theochem) 1997, V389, P37 ZCAPLUS
(10) Chiu, Y; Phys Rev B 1997, V55, P6022 ZCAPLUS
(11) Cotton, F; Chemical Application of Group Theory 1990
(12) Ganelin, P; Fullerenes, Recent Advances in the Chemistry and Physics
    of Fullerenes and Related Materials 1994
```

(13) Kato, T; J Phys Chem 1997, V97, P13425

- (14) Kroto, H; Nature 1987, V329, P529 ZCAPLUS
- (15) Kroto, H; Nature 1995, V318, P162
- (16) Manalopoulos, D; Chem Phys Lett 1991, V187, P11
- (17) Merritt, C; PhD thesis, The Catholic University of America 1998
- (18) Moro, L; J Phys Chem 1997, V97, P6801
- (19) Reeves, M; Phys Rev B 1993, V47, P6065 ZCAPLUS
- (20) Sarkas, H; J Phys Chem 1996, V100, P5169 ZCAPLUS
- (21) Schmalz, T; J Am Chem Soc 1988, V110, P113
- IT 309242-98-6, [4,5,6] Fullerene-C30-C2v

(geminals and Schlegel diagrams of fullerenes and metallofullerenes)

- RN 309242-98-6 ZCAPLUS
- CN [4,5,6]Fullerene-C30-C2v (9CI) (CA INDEX NAME)



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STRUCTURE FILE UPDATES: 28 FEB 2005 HIGHEST RN 839671-97-5 DICTIONARY FILE UPDATES: 28 FEB 2005 HIGHEST RN 839671-97-5

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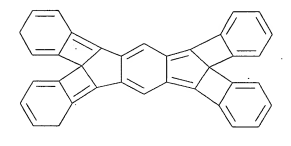
- L11 ANSWER 1 OF 2 REGISTRY COPYRIGHT 2005 ACS on STN
- RN 334833-14-6 REGISTRY
- ED Entered STN: 07 May 2001
- CN 1H,6H-Tetrakisbenzo[3,4]cyclobut[1,2-a:1',2'-b:1'',2''-g:1''',2'''-h]-s-indacene (9CI) (CA INDEX NAME)
- FS 3D CONCORD
- MF C36 H20



SR CA Index Guide or Ring Systems Handbook

## Ring System Data

Elemental	Elemental	Size of	Ring System	Ring	RID
Analysis	Sequence	the Rings	Formula	Identifier	Occurrence
EA	l ES	SZ	RF	RID	Count
=============	+==========	+= <b>==</b> =====	+=========	+=======	-=======
C4-C4-C4-C4-	C4-C4-C4-C4-	4-4-4-4-5-5-	JC36	86620.1.2	1
C5-C5-C6-C6-	C5-C5-C6-C6-	6-6-6-6-6	1		
C6-C6-C6	C6-C6-C6				



This compound is not actually cited in any abstracts or citations. It just was requitered by a company and assigned a registry number and index name by Chemical abstracts.)

# Calculated Properties (CALC)

PROPERTY (CODE)	VALUE	CONDITION  NOTE	
Bioconc. Factor (BCF)	503742		CD
Bioconc. Factor (BCF)	1503742	1 * 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	CD
Bioconc. Factor (BCF)	1503742		CD
Bioconc. Factor (BCF)	503742		CD
Bioconc. Factor (BCF)	503742	pH 10   (1) A	CD
H acceptors (HAC)	0	(1) A	CD
H donors (HD)	10	(1) A	CD
Koc (KOC)	419964	pH 1   (1) A	CD
Koc (KOC)	419964	pH 4	$^{\prime}$ CD
Koc (KOC)	419964	pH 7   (1) A	$^{\prime}$ CD
Koc (KOC)	1419964	pH 8   (1) A	$^{\prime}$ CD
Koc (KOC)	419964	pH 10   (1) A	$^{\prime}$ CD
logD (LOGD)	7.81	pH 1   (1) A	$^{\prime}$ CD
logD (LOGD)	7.81	pH 4   (1) A	$^{\prime}$ CD
logD (LOGD)	7.81	pH 7  (1) A	$^{\prime}$ CD
logD (LOGD)	7.81	pH 8   (1) A	'CD

```
logD (LOGD)
                           17.81
                                          |pH 10
                                                    (1) ACD
                           |7.806+/-0.618|
logP (LOGP)
                                                    |(1) ACD
Molar Solubility (SLB.MOL) | < 0.01 mol/L
                                         |pH 1
                                                    |(1) ACD
Molar Solubility (SLB.MOL) | < 0.01 mol/L
                                                    |(1) ACD
                                         |pH 4
Molar Solubility (SLB.MOL) |<0.01 mol/L |pH 7
                                                    (1) ACD
Molar Solubility (SLB.MOL) | < 0.01 mol/L
                                         8 Hq|
                                                    |(1) ACD
Molar Solubility (SLB.MOL) | < 0.01 mol/L
                                          |pH 10
                                                    |(1) ACD
Molecular Weight (MW) | 452.54
                                                    |(1) ACD
```

2 63 6

(1) Calculated using Advanced Chemistry Development (ACD/Labs) Software Solaris V4.67 ((C) 1994-2005 ACD/Labs)

See HELP PROPERTIES for information about property data sources in REGISTRY.